TCEQ Interoffice Memorandum

To: Tony Walker

Director, TCEQ Region 4, Dallas/Fort Worth

Alyssa Taylor

Air Section Manager, TCEQ Region 4, Dallas/Fort Worth

From: Shannon Ethridge, M.S. S.E.

Toxicology Division, Chief Engineer's Office

Date: June 22, 2011

Subject: Toxicological Evaluation of Results from Ambient Air Samples for Volatile

Organic Compounds Collected Upwind (Latitude 33.37959, Longitude -97.37039) and Downwind (Latitude 33.38111, Longitude -97.36943) of the Burlington

Resources - McMurrey Ranch 22D, 28H, 25H, 35H Site Near Sanger, Denton

County, Texas

Samples Collected on April 18, 2011, ACLs 1104052 and 1104047 (Lab Samples

1104052-001 and 1104047-001)

Key Points

- The reported concentration of 2-methylpentane in the downwind sample (Lab Sample 1104047-001) exceeded its odor air monitoring comparison value (AMCV) (Table 1). The reported level of this chemical would be expected to cause an odor if exposure were to occur, which is consistent with the TCEQ regional staff report of a chemical odor during the sampling event.
- At this time, the general public would not be expected to be exposed to emissions from this facility due to the location of the nearest residence (approximately 3,500 feet to the south of the site). However, if land use around the facility changes in the future, exposure may be a possibility.

Background

On April 18, 2011, Texas Commission on Environmental Quality (TCEQ) Region 4 Air Investigators collected two 30-minute canister samples, upwind (Lab Sample 1104052-001, Latitude 33.37959, Longitude -97.37039) and downwind (Lab Sample 1104047-001, Latitude 33.38111, Longitude -97.36943) of the Burlington Resources - McMurrey Ranch 22D, 28H, 25H, 35H Site near Sanger, Denton County, Texas. The samples were collected as a follow-up to a previous investigation. The investigators noted a chemical odor during the upwind and downwind sampling events. Meteorological conditions measured at the site or nearest stationary

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ambient air monitoring site indicated that the ambient temperature was 89.7°F with a relative humidity of 47.6%. Winds were from the south (180°) at 10.8 miles per hour at the downwind sampling location. Winds were from the south (180°) at 6.4 miles per hour at the upwind sampling location. The sampling site for the downwind sample was approximately 100 to 300 feet from the possible source. The nearest residential property is located approximately 3,500 feet to the south of the possible source. The samples were sent to the TCEQ laboratory in Austin, Texas, and analyzed for a range of volatile organic compounds (VOCs). The list of the target analytes that were evaluated in this review are provided in Attachment A. The VOC concentrations were reported in parts per billion by volume (ppb_v) (Attachment B and Tables 2 and 3). Please note that the available canister technology and analysis method can not capture and/or analyze for all chemicals.

Results and Evaluation

Reported VOC concentrations were compared to TCEQ's short-term health- and/or welfare-based AMCVs (Tables 2 and 3). Short-term AMCVs are guidelines used to evaluate ambient concentrations of a chemical in air and to determine its potential to result in adverse health effects, adverse vegetative effects, or odors. Health AMCVs are set to provide a margin of safety, and are set well below levels at which adverse health effects are reported in the scientific literature. If a chemical concentration in ambient air is less than its comparison value, no adverse health effects are expected to occur. If a chemical concentration exceeds its comparison value it does not necessarily mean that adverse effects will occur, but rather that further evaluation is warranted.

Eighty-three of the 84 VOCs monitored in downwind sample and all 84 VOCs monitored in the upwind sample were either not detected or were detected below their respective short-term AMCVs. Exposure to levels of these VOCs measured in these samples would not be expected to cause short-term adverse health effects, adverse vegetative effects, or odors.

The reported concentration 2-methylpentane exceeded its odor AMCV (Table 1). The reported level of this chemical would be expected to cause an odor if exposure were to occur, which is consistent with the TCEQ regional staff report of a chemical odor during the sampling event.

At this time, the general public would not be expected to be exposed to emissions from this facility due to the location of the nearest residence (approximately 3,500 feet to the south of the site). However, if land use around the facility changes in the future, exposure may be a possibility.

Please call me at (512) 239-1822 if you have any questions regarding this evaluation.

Table 1. Exceedance in Lab Sample 1104047-001

Chemical	Measured Concentration (ppb _v)	Short-term health AMCV (ppb _v)	Short-term odor AMCV (ppb _v)	Does it exceed the short-term, health AMCV?	Does it exceed the short-term, odor AMCV ?
2-methylpentane	97¹	1,000	83	No	Yes

 1 D2 - Sample concentration was calculated using a dilution factor of 80.00 and the diluted sample was analyzed on 5/4/2011.

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Attachment A

List of Target Analytes for Canister Samples

ethane ethylene acetylene propane propylene dichlorodifluoromethane methyl chloride isobutane vinyl chloride 1-butene 1.3-butadiene n-butane t-2-butene bromomethane c-2-butene 3-methyl-1-butene

isopentane

trichlorofluoromethane

1-pentene n-pentane isoprene t-2-pentene

1,1-dichloroethylene

c-2-pentene

methylene chloride 2-methyl-2-butene 2,2-dimethylbutane cyclopentene

4-methyl-1-pentene 1.1-dichloroethane cyclopentane 2,3-dimethylbutane 2-methylpentane 3-methylpentane

2-methyl-1-pentene + 1-hexene

n-hexane chloroform t-2-hexene c-2-hexene

1,2-dichloroethane methylcyclopentane 2,4-dimethylpentane 1,1,1-trichloroethane

benzene

carbon tetrachloride

cyclohexane 2-methylhexane 2,3-dimethylpentane 3-methylhexane 1,2-dichloropropane trichloroethylene 2,2,4-trimethylpentane

2-chloropentane

n-heptane

c-1,3-dichloropropylene methylcyclohexane

t-1,3-dichloropropylene 1.1.2-trichloroethane 2,3,4-trimethylpentane toluene

2-methylheptane 3-methylheptane 1.2-dibromoethane

n-octane

tetrachloroethylene chlorobenzene ethylbenzene m & p-xylene styrene

1,1,2,2-tetrachloroethane

o-xylene n-nonane

isopropylbenzene n-propylbenzene m-ethyltoluene p-ethyltoluene

1,3,5-trimethylbenzene

o-ethyltoluene

1,2,4-trimethylbenzene

n-decane

1,2,3-trimethylbenzene m-diethylbenzene p-diethylbenzene n-undecane

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Attachment B

5/6/2011

Texas Commission on Environmental Quality

Laboratory and Quality Assurance Section P.O. Box 13087, MC-165 Austin, Texas 78711-3087 (512) 239-1716

Laboratory Analysis Results ACL Number: 1104047

ACL Lead: Karen Bachtel

Region: T04

Date Received: 4/20/2011

Project(s): Barnett Shale

Facility(ies) Sampled	City	County	Facility Type
Burlington Resources	Sanger .	Denton	

Laboratory Procedure(s) Performed:

Analysis: AMOR006

Determination of VOC Canisters by GC/MS Using Modified Method TO-15

Prior to analysis, subatmospheric samples are pressurized to twice the collected volume using a sample dilution system. For analysis, a known volume of a sample is directed from the canister into a multitrap cryogenic concentrator. Internal standards are added to the sample stream prior to the trap. The concentrated sample is thermally desorbed and carried onto a GC column for separation. The analytical strategy involves using a GC with dual columns that are coupled to a mass selective detector (MSD) and a flame ionization detector (FID). Mass spectra for individual peaks in the total ion chromatogram are then used for target compound identification and quantitation. The fragmentation pattern is compared with stored spectra taken under similar conditions in order to identify the compound. For any given compound, the intensity of the quantitation fragment is compared with the system response to the fragment for known amounts of the compound. This establishes the compound concentration in the sample. For non-target compound peaks which are at least one-half the height of the internal standard, a library search is performed in an attempt to identify the compound solely upon fracture patterns. These tentatively identified compounds (TIC's) are reported as a sample specific footnote. Accurate quantitation of TICs is not possible. The FID is used for the quantitation of ethane, ethylene, acetylene, propylene and propane and identification is based on matching retention times of standards containing known analytes.

Sample(s) Received

Field ID Number: 01178 Sampling Site: McMurrey Ranch 22D Laboratory Sample Number: 1104047-001

Sampled by: Abayomi Dedeke

Date & Time Sampled: 04/18/11 11:50:00 Valid Sample: Yes

Comments:

Canister 01178 was used to collect a 30-minute sample using CO-010.

Please note that this analytical technique is not capable of measuring all compounds which might have adverse health effects. For questions on the analytical procedures please contact the laboratory manager at (512) 239-4894. For an update on the health effects evaluation of these data, please contact the Toxicology Division at (512) 239-1795.

Reviewed By

Technical Specialist:

David Manis

Laboratory Analysis Results

ACL Number: 1104047 Analysis Code: AMOR006

Note: Results are reported in units of parts per billion by volume (ppbv) 1104047-001 Lab ID 01178 Field ID Canister ID 01178 Analysis Date 04/28/11 LOD Concentration SDL Flags** Concentration SDL Flags** Compound ethane 0.50 5100 1.0 D2,T ethylene 0.50 ND 1.0 D1,T 0.31 1.0 J,D1,T 0.50 acetylene propane 0.50 6300 1.0 D2,T0.50 0.44 1.0 J,D1,T propylene dichlorodifluoromethane 0.20 0.63 0.40 L,D1 methyl chloride 0.20 0.88 0.40 L,D1 isobutane 0.23 1000 0.46 D20.34 D1 vinyl chloride 0.17 ND 0.40 1-butene 0.20 0.36 J,D1 1,3-butadiene 0.27 ND 0.54 D1 n-butane 0.20 2400 0.40 D20.18 ИD 0.36 DI t-2-butene bromomethane 0.27 ND 0.54 D1 c-2-butene 0.27 ND 0.54 Di 3-methyl-1-butene 0.23 0.13 0.46 J.DI 0.54 isopentane 0.27 600 D2 trichlorofluoromethane 0.29 0.32 0.58 J,D1 0.54 0.27 ND D1 1-pentene 0.54 n-pentane 0.27 630 D20.27 ND 0.54 Ď1 isoprene t-2-pentene 0.27 ND 0.54 Dl D1 0.18 ND 0.36 1,1-dichloroethylene 0.25 ND 0.50 Dl c-2-pentene 0.28 Dl methylene chloride 0.14 ND 2-methyl-2-butene 0.23 ND 0.46 D1 2,2-dimethylbutane 0.21 4.2 0.42 DI cyclopentene 0.20 ND 0.40 DI 0.44 0.22 ND D1 4-methyl-1-pentene 1,1-dichloroethane 0.19 ND 0.38 D1 22 0.27 0.54 D2cyclopentane 2,3-dimethylbutane 0.28 12 0.56 D1 2-methylpentane 0.27 97 0.54 D2 3-methylpentane 0.2354 0.46 D22-methyl-1-pentene + 1-hexene 0.20 0.30 0.40 J,DI n-hexane 0.20120 0.40 D2chloroform 0.21 ND 0.42DI 0.27 ND0.54 D1 o-2-hexene 0.27 ND 0.54 D1 1,2-dichloroethane 0.27 ND 0.54 D1 0.27 40 0.54 D2 methylcyclopentane 2,4-dimethylpentane 0.27 2.6 0.54 D1 1,1,1-trichloroethane 0.26 ND 0.52 DI 0.27 18 0.54 D1 carbon tetrachloride 0.27 0.31 0.54 J,D1 cyclohexane 0.24 33 0.48 D2 2-methylhexane 0.27 19 0.54 DI 2,3-dimethylpentane 0.26 5.5 0.52 D1

Laboratory Analysis Results

ACL Number: 1104047 Analysis Code: AMOR006

Lab ID		110	4047-001				
Compound	LOD	Concentration	SDL	Flags**	Concentration	SDL	Flags**
3-methylhexane	0.20	17	0.40	D2			
1,2-dichloropropane	0.17	ND	0.34	D1			
trichloroethylene	0.29	ND	0.58	D1			
2,2,4-trimethylpentane	0.24	ND	0.48	D1			
2-chioropentane	0.27	ND	0.54	D1			
n-heptanc	0.25	27 .	0.50	D2			
c-1,3-dichloropropylene	0.20	ND	0.40	DI			
methyloyclohexane	0.26	25	0.52	D2			
-1,3-dichloropropylene	0.20	ND	0.40	DI			
1,1,2-trichloroethane	0.21	ND	0.42	D1 ·			
2,3,4-trimethylpentane	0.24	ND	0.48	D1			
toluene	0.27	20	0.54	D1			
2-methylheptane	0.20	6.5	0.40	. D1			
3-methylheptane	0.23	4.1	0.46	·Dl			
,2-dibromoethane	0.20	0.22	0.40	J,D1			
n-octane	0.19	11	0.38	D1			
etrachlomethylene	0.24	0.16	0.48	J,D1			
chlorobenzene	0.27	ND	0.54	DI			
sthylbenzene	0.27	1.5	0.54	Dl			
n & p-xylene	0.27	7.6	0.54	DI			
styrene	0.27	ND	0.54	D1			
1,1,2,2-tetrachloroethane	0.20	ND	0.40	D1			
-xylene	0.27	2.5	0.54	D1			
1-nonane	0.22	2.9	0.44	D1 .			
sopropylbenzene	0.24	ND	0.48	D1			
1-propylbenzene	0.27	ND	0.54	D1			
n-ethyltoluene	0.11	0.67	0.22	L,D1			
-ethyltoluene	0.16	0.36	0.32	L,DI			
,3,5-trimethylbenzene	0.25	ND	0.50	Dl			
-ethyltoluene	0.13	0.32	0.26	L ₂ D1			
,2,4-trimethylbenzene	0.27	1.1	0.54	L,DI			
-decane	0.27	0.86	0.54	L,DI			
,2,3-trimethylbenzene	0.27	ND	0.54	. D1			
n-diethylbenzene	0.27	. ND	0.54	D1			
-diethylbenzene	0.27	0.29	0.54	J,D1			
-undecane	0.27	ND	0.54	Di			

Laboratory Analysis Results ACL Number: 1104047

Analysis Code: AMOR006

Note: Results are reported in units of parts per billion by volume (ppbv)

LOD - Limit of Detection.

ND - not detected

NQ - concentration can not be quantified.

SDL - Sample Detection Limit (LOD adjusted for dilutions).

INV - Invalid.

J - Reported concentration is below SDL.

L - Reported concentration is at or above the SDL and is below the lower limit of quantitation.

E - Reported concentration exceeds the upper limit of instrument calibration.

M - Result modified from previous result.

T- Data was not confirmed by a confirmational analysis. Data is tentatively identified.

* SDL is equal to LOD

** Quality control flags explanations are listed on the last page of this report.

TCEQ laboratory customer support may be reached at kbachtel@tceq.state.tx.us

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Laboratory Analysis Results ACL Number: 1104047 Analysis Code: AMOR006

Quality Control Notes:

D1 -Sample concentration was calculated using a dilution factor of 4.00. D2 -Sample concentration was calculated using a dilution factor of 80.00 and the diluted sample was analyzed on 05/04/2011.

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5/5/2011

Texas Commission on Environmental Quality

Laboratory and Quality Assurance Section P.O. Box 13087, MC-165 Austin, Texas 78711-3087 (512) 239-1716

Laboratory Analysis Results ACL Number: 1104052

ACL Lead: Karen Bachtel

Region: T04

Date Received: 4/21/2011

Project(s): Barnett Shale

Facility(ies) Sampled	City	County	Facility Type
Burlington Resources	Sanger	Denton	

Laboratory Procedure(s) Performed:

Analysis: AMOR006

Determination of VOC Canisters by GC/MS Using Modified Method TO-15

Procedure:

Prior to analysis, subatmospheric samples are pressurized to twice the collected volume using a sample dilution system. For analysis, a known volume of a sample is directed from the canister into a multitrap cryogenic concentrator. Internal standards are added to the sample stream prior to the trap. The concentrated sample is thermally desorbed and carried onto a GC column for separation. The analytical strategy involves using a GC with dual columns that are coupled to a mass selective detector (MSD) and a flame ionization detector (FID). Mass spectra for individual peaks in the total ion chromatogram are then used for target compound identification and quantitation. The fragmentation pattern is compared with stored spectra taken under similar conditions in order to identify the compound. For any given compound, the intensity of the quantitation fragment is compared with the system response to the fragment for known amounts of the compound. This establishes the compound concentration in the sample. For non-target compound peaks which are at least one-half the height of the internal standard, a library search is performed in an attempt to identify the compound solely upon fracture patterns. These tentatively identified compounds (TIC's) are reported as a sample specific footnote. Accurate quantitation of TICs is not possible. The FID is used for the quantitation of ethane, ethylene, acetylene, propylene and propane and identification is based on matching retention times of standards containing known analytes.

Sample(s) Received

Field ID Number: 20294

Laboratory Sample Number: 1104052-001

Sampled by: Daniel Atambo

Sampling Site: McMurrey Ranch

Date & Time Sampled: 04/18/11 11:50:00 Valid Sample: Yes

Comments:

Canister 20294 was used to collect a 30-minute sample using OFC-022.

Please note that this analytical technique is not capable of measuring all compounds which might have adverse health effects. For questions on the analytical procedures please contact the laboratory manager at (512) 239-4894. For an update on the health effects evaluation of these data, please contact the Toxicology Division at (512) 239-1795.

Analyst: Jaydeep Patel

Reviewed By: Saren Bachtel

Technical Specialist: David Manis

Laboratory Analysis Results ACL Number: 1104052

Analysis Code: AMOR006

Leb ID			04052-001				
Leb ID							
Field ID			20294	-			
Canister ID		20294					
Analysis Date			04/28/11			,	
Compound	LOD	Concentration	SDL	Flags**	Concentration	SDL	Flags**
ethane	0.50	6.1	1.0	D1,T			
ethylene	0.50	0.74	1.0	J,D1,T			ļ
acetylene	0.50	0.38	. 1.0	J,D1,T			
propane	0.50	2.5	1.0	DI,T			
propylene	0.50	0.37	1.0	J,D1,T			<u> </u>
dichlorodifluoromethane	0.20	0.50	0.40	L,DI			
methyl chloride	0.20	0.82	0.40	L,DI			
sobutane	0.23	0.87	0.46	L _i D1			
vinyl chloride	0.17	ND .	0.34	DI			<u> </u>
I-butene	0.20	0.37	0.40	J,DI			
1,3-butadiene	0.27	0.13	0.54	J,D1			
n-butane	0.20	1.4	0.40	L,D1			
-2-butene	0.18	ND	0.36	D1			
bromomethane	0.27	ND	0.54	D1 ·			
>2-butene	0.27	ND	0.54	D1			
3-methyl-1-butene	0.23	ND	0.46	D1			
sopentane	0.27	0.67	0.54	L ₂ D1			
richlorofluoromethane	0.29	0.36	0.58	J,D1			
-pentene	0.27	ND	0.54	DI			
s-pentane	0.27	ND	0.54	D1			
soprene	- 0.27	ND	0.54	DI			
-2-pentene	0.27	ND	0.54	DI			
,1-dichloroethylene	0.18	ND	0.36	D1			
-2-pentene	0.25	ND	0.50	. D1			
nethylene chloride	0.14	ND	0.28	D1			
2-methyl-2-butene	0.23	ND	0.46	D1			
2,2-dimethylbutane	0.21	ND	0.42	D1			
cyclopentene	0.20	ND	0.40	Ď1			
i-methyl-1-pentene	0.22	ND	0.44	D1			
,1-dichloroethane	0.19	ND	0.38	DI			
cyclopentane	0.27	0.13	0.54	J.DÍ			<u> </u>
2,3-dimethy/butane	0.28	ND	0.56	DI			
2-methylpentane	0.27	ND	0.54	D1			
	0.27	ND	0.46	DI			
-methylpentane	0.20	ND	0.40	DI			
l-methyl-1-pentene + 1-hexene	0.20	ND	0.40	DI	· ·		
hiexane	0.20	0.13					
hloroform			0.42	J,D1			
-2-hexane	0.27	ND	0.54	DI			
-2-hexene	0.27	ND	0.54	DI .			
,2-dichloroethane	0.27	ND	0.54	D1			
nethylcyclopentane	0.27	ND	0.54	DI.			
,4-dimethylpentane	0.27	ND	0.54	D1			
,1,1-trichloroethane	0.26	ND	0.52	Dl			
enzene	0.27	0.31	0.54	J,D1			
arbon tetrachloride	0.27	0.32	0.54	J,D1			
yclohexane	0.24	ND	0.48	D1			
-methylhexane	0.27	ND	0.54	D1		- 1	

Laboratory Analysis Results

ACL Number: 1104052 Analysis Code: AMOR006

Lab ID		110	4052-001				
Compound	LOD	Concentration	SDL	Flags**	Concentration	SDL	Flags**
3-methylhexane	0.20	ND	0.40	D1			
I,2-dichloropropane	0.17	ND	0.34	Dl			
trichloroethylene	0.29	ND	0.58	DI			
2,2,4-trimethylpentane	0.24	ND	0.48	D1			
2-chloropentane	0.27	ND	0.54	. D1			
n-heptane	0.25	ND	0.50	Dl			
c-1,3-dichloropropylene	0.20	ND	0.40	D1			
methylcyclohexane	0.26	ND	0.52	DI			
t-1,3-dichloropropylene	0.20	ND	0.40	D1		i	
1,1,2-trichloroethane	0.21	ND	0.42	DI			
2,3,4-trimethy(pentane	0.24	ND	0.48	DI			
oluene	0.27	0.25	0.54	J,DI			
2-methylheptane	0.20	ND	0.40	D1			
3-methylheptane	0.23	ND	0.46	D1			
,2-dibromoethane	0.20	ИD	0.40	D1			
1-octane	0.19	ND	0.38	D1			
tetrachloroethylene	0.24	0.12	0.48	J,D1			
chlorobenzene	0.27	ND	0.54	Dl			
sthylbenzene	0.27	ND	0.54	DI			
n & p-xylene	0.27	0.43	0.54	J,D1			
tyrene	0.27	ND	0.54	DI			
,1,2,2-tetrachloroethane	. 0.20	ND.	0.40	DI :			
o-xylene	0.27	ND	0.54	D1			
n-nonane	0.22	ND	0.44	D1			
sopropylbenzene	0.24	ND	0.48	D1			
-propylbenzene	0.27	ND	0.54	D1			
n-ethyltoluene	0.11	ND	0.22	Dl			
-ethyltolucue	0.16	ND	0.32	DI			
,3,5-trimethylbenzene	0.25	ND	0.50	D1			
-ethyltoluene	0.13	ND	0.26	DI			
,2,4-trimethylbenzene	0.27	ND	0.54	D1			
-decane	0.27	ND	0.54	DI			
,2,3-trimethylbenzene	0,27	ND	0.54	DI			-
n-diethylbenzene	0.27	ND	0.54	Dl			
-diethylbenzene	0.27	0.14	0.54	J,D1			
-undecane	0.27	ND	0.54	D1			

Laboratory Analysis Results ACL Number: 1104052

Analysis Code: AMOR006

Note: Results are reported in units of parts per billion by volume (ppbv)

LOD - Limit of Detection.

ND - not detected

NQ - concentration can not be quantified.

SDL - Sample Detection Limit (LOD adjusted for dilutions).

INV - Invalid.

J - Reported concentration is below SDL.

L - Reported concentration is at or above the SDL and is below the lower limit of quantitation.

E - Reported concentration exceeds the upper limit of instrument calibration.

M - Result modified from previous result.

T- Data was not confirmed by a confirmational analysis. Data is tentatively identified.

* SDL is equal to LOD

** Quality control flags explanations are listed on the last page of this report.

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Laboratory Analysis Results ACL Number: 1104052 Analysis Code: AMOR006

Quality Control Notes:

D1 -Sample concentration was calculated using a dilution factor of 4.02.

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Table 2. Comparison of Monitored Concentrations in Lab Sample 1104047-001 to TCEQ Short-Term AMCVs

Lab Sample ID	1104047-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
1,1,1-Trichloroethane	380,000	1,700	0.26	ND	D1	0.52
1,1,2,2-Tetrachloroethane	7,300	10	0.2	ND	D1	0.4
1,1,2-Trichloroethane	Not Available	100	0.21	ND	D1	0.42
1,1-Dichloroethane	110,000	1,000	0.19	ND	D1	0.38
1,1-Dichloroethylene	Not Available	180	0.18	ND	D1	0.36
1,2,3-Trimethylbenzene	Not Available	250	0.27	ND	D1	0.54
1,2,4-Trimethylbenzene	Not Available	250	0.27	1.1	L,D1	0.54
1,2-Dibromoethane	10,000	0.5	0.2	0.22	J,D1	0.4
1,2-Dichloroethane	6,000	40	0.27	ND	D1	0.54
1,2-Dichloropropane	250	100	0.17	ND	D1	0.34
1,3,5-Trimethylbenzene	Not Available	250	0.25	ND	D1	0.5
1,3-Butadiene	230	1,700	0.27	ND	D1	0.54
1-Butene	360	50,000	0.2	0.36	J,D1	0.4
1-Pentene	100	2,600	0.27	ND	D1	0.54
2,2,4-Trimethylpentane	Not Available	750	0.24	ND	D1	0.48
2,2-Dimethylbutane (Neohexane)	Not Available	1,000	0.21	4.2	D1	0.42
2,3,4-Trimethylpentane	Not Available	750	0.24	ND	D1	0.48
2,3-Dimethylbutane	Not Available	990	0.28	12	D1	0.56
2,3-Dimethylpentane	Not Available	850	0.26	5.5	D1	0.52
2,4-Dimethylpentane	290,000	850	0.27	2.6	D1	0.54
2-Chloropentane (as chloroethane)	Not Available	190	0.27	ND	D1	0.54
2-Methyl-1-Pentene +1-Hexene	20	500	0.2	0.3	J,D1	0.4
2-Methyl-2-Butene	250	500	0.23	ND	D1	0.46
2-Methylheptane	Not Available	750	0.2	6.5	D1	0.4

Lab Sample ID	1104047-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
2-Methylhexane	Not Available	750	0.27	19	D1	0.54
2-Methylpentane (Isohexane)	83	1,000	0.27	97	D2	0.54
3-Methyl-1-Butene	250	8,000	0.23	0.13	J,D1	0.46
3-Methylheptane	Not Available	750	0.23	4.1	D1	0.46
3-Methylhexane	Not Available	750	0.2	17	D2	0.4
3-Methylpentane	Not Available	1,000	0.23	54	D2	0.46
4-Methyl-1-Pentene (as hexene)	20	500	0.22	ND	D1	0.44
Acetylene	620,000	25,000	0.5	0.31	J,D1,T	1
Benzene	2,700	180	0.27	18	D1	0.54
Bromomethane (methyl bromide)	21,000	30	0.27	ND	D1	0.54
c-1,3-Dichloropropylene	Not Available	10	0.2	ND	D1	0.4
c-2-Butene	2,100	15,000	0.27	ND	D1	0.54
c-2-Hexene	Not Available	500	0.27	ND	D1	0.54
c-2-Pentene	Not Available	2,600	0.25	ND	D1	0.5
Carbon Tetrachloride	97,000	20	0.27	0.31	J,D1	0.54
Chlorobenzene (phenyl chloride)	210	100	0.27	ND	D1	0.54
Chloroform (trichloromethane)	85,000	20	0.21	ND	D1	0.42
Cyclohexane	420	1,000	0.24	33	D2	0.48
Cyclopentane	Not Available	1,200	0.27	22	D2	0.54
Cyclopentene	Not Available	2,900	0.2	ND	D1	0.4
Dichlorodifluoromethane	Not Available	10,000	0.2	0.63	L,D1	0.4
Ethane	180,000	Simple Asphyxiant*	0.5	5,100	D2,T	1
Ethylbenzene	170	20,000	0.27	1.5	D1	0.54
Ethylene	270,000	500,000	0.5	ND	D1,T	1
Isobutane	2,040	8,000	0.23	1,000	D2	0.46
Isopentane (2-methylbutane)	1,300	1,200	0.27	600	D2	0.54

Lab Sample ID	1104047-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
Isoprene	5	20	0.27	ND	D1	0.54
Isopropylbenzene (cumene)	100	500	0.24	ND	D1	0.48
m & p-Xylene (as mixed isomers)	80	1,700	0.27	7.6	D1	0.54
m-Diethylbenzene	70	460	0.27	ND	D1	0.54
Methyl Chloride (chloromethane)	Not Available	500	0.2	0.88	L,D1	0.4
Methylcyclohexane	150	4,000	0.26	25	D2	0.52
Methylcyclopentane	1,700	750	0.27	40	D2	0.54
Methylene Chloride (dichloromethane)	160,000	3,500	0.14	ND	D1	0.28
m-Ethyltoluene	18	250	0.11	0.67	L,D1	0.22
n-Butane	1,200,000	8,000	0.2	2,400	D2	0.4
n-Decane	620	1,750	0.27	0.86	L,D1	0.54
n-Heptane	670	850	0.25	27	D2	0.5
n-Hexane	1,500	1,800	0.2	120	D2	0.4
n-Nonane	2,200	2,000	0.22	2.9	D1	0.44
n-Octane	1,700	750	0.19	11	D1	0.38
n-Pentane	1,400	1,200	0.27	630	D2	0.54
n-Propylbenzene	3.8	250	0.27	ND	D1	0.54
n-Undecane	Not Available	550	0.27	ND	D1	0.54
o-Ethyltoluene	Not Available	250	0.13	0.32	L,D1	0.26
o-Xylene	380	1,700	0.27	2.5	D1	0.54
p-Diethylbenzene	0.39	460	0.27	0.29	J,D1	0.54
p-Ethyltoluene	8.3	250	0.16	0.36	L,D1	0.32
Propane	1,500,000	Simple Asphyxiant*	0.5	6300	D2,T	1
Propylene	13,000	Simple Asphyxiant*	0.5	0.44	J,D1,T	1
Styrene	25	5,100	0.27	ND	D1	0.54
t-1,3-Dichloropropylene	Not Available	10	0.2	ND	D1	0.4

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Lab Sample ID	1104047-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
t-2-Butene	2,100	15,000	0.18	ND	D1	0.36
t-2-Hexene	Not Available	500	0.27	ND	D1	0.54
t-2-Pentene	Not Available	2,600	0.27	ND	D1	0.54
Tetrachloroethylene	770	1,000	0.24	0.16	J,D1	0.48
Toluene	170	4,000	0.27	20	D1	0.54
Trichloroethylene	3,900	100	0.29	ND	D1	0.58
Trichlorofluoromethane	5,000	10,000	0.29	0.32	J,D1	0.58
Vinyl Chloride	Not Available	26,000	0.17	ND	D1	0.34

^{*}A simple asphyxiant displaces air, lowering the partial pressure of oxygen and causing hypoxia at sufficiently high concentrations. ppbv - parts per billion by volume

ND - Not Detected.

NQ - Concentration can not be quantified.

LOD - Limit of Detection.

SDL - Sample Detection Limit (LOD adjusted for dilutions).

INV - Invalid.

- J Reported concentration is below SDL.
- L Reported concentration is at or above the SDL and is below the lower limit of quantitation.
- E Reported concentration exceeds the upper limit of instrument calibration.
- M Result modified from previous result.
- T Data was not confirmed by a confirmational analysis. Data is tentatively identified.
- D1 Sample concentration was calculated using a dilution factor of 4.00.
- D2 Sample concentration was calculated using a dilution factor of 80.00 and the diluted sample was analyzed on 5/4/2011.

Table 3. Comparison of Monitored Concentrations in Lab Sample 1104052-001 to TCEQ Short-Term AMCVs

Lab Sample ID	1104052-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
1,1,1-Trichloroethane	380,000	1,700	0.26	ND	D1	0.52
1,1,2,2-Tetrachloroethane	7,300	10	0.2	ND	D1	0.4
1,1,2-Trichloroethane	Not Available	100	0.21	ND	D1	0.42
1,1-Dichloroethane	110,000	1,000	0.19	ND	D1	0.38
1,1-Dichloroethylene	Not Available	180	0.18	ND	D1	0.36
1,2,3-Trimethylbenzene	Not Available	250	0.27	ND	D1	0.54
1,2,4-Trimethylbenzene	Not Available	250	0.27	ND	D1	0.54
1,2-Dibromoethane	10,000	0.5	0.2	ND	D1	0.4
1,2-Dichloroethane	6,000	40	0.27	ND	D1	0.54
1,2-Dichloropropane	250	100	0.17	ND	D1	0.34
1,3,5-Trimethylbenzene	Not Available	250	0.25	ND	D1	0.5
1,3-Butadiene	230	1,700	0.27	0.13	J,D1	0.54
1-Butene	360	50,000	0.2	0.37	J,D1	0.4
1-Pentene	100	2,600	0.27	ND	D1	0.54
2,2,4-Trimethylpentane	Not Available	750	0.24	ND	D1	0.48
2,2-Dimethylbutane (Neohexane)	Not Available	1,000	0.21	ND	D1	0.42
2,3,4-Trimethylpentane	Not Available	750	0.24	ND	D1	0.48
2,3-Dimethylbutane	Not Available	990	0.28	ND	D1	0.56
2,3-Dimethylpentane	Not Available	850	0.26	ND	D1	0.52
2,4-Dimethylpentane	290,000	850	0.27	ND	D1	0.54
2-Chloropentane (as chloroethane)	Not Available	190	0.27	ND	D1	0.54
2-Methyl-1-Pentene +1-Hexene	20	500	0.2	ND	D1	0.4
2-Methyl-2-Butene	250	500	0.23	ND	D1	0.46
2-Methylheptane	Not Available	750	0.2	ND	D1	0.4

Lab Sample ID	1104052-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
2-Methylhexane	Not Available	750	0.27	ND	D1	0.54
2-Methylpentane (Isohexane)	83	1,000	0.27	ND	D1	0.54
3-Methyl-1-Butene	250	8,000	0.23	ND	D1	0.46
3-Methylheptane	Not Available	750	0.23	ND	D1	0.46
3-Methylhexane	Not Available	750	0.2	ND	D1	0.4
3-Methylpentane	Not Available	1,000	0.23	ND	D1	0.46
4-Methyl-1-Pentene (as hexene)	20	500	0.22	ND	D1	0.44
Acetylene	620,000	25,000	0.5	0.38	J,D1,T	1
Benzene	2,700	180	0.27	0.31	J,D1	0.54
Bromomethane (methyl bromide)	21,000	30	0.27	ND	D1	0.54
c-1,3-Dichloropropylene	Not Available	10	0.2	ND	D1	0.4
c-2-Butene	2,100	15,000	0.27	ND	D1	0.54
c-2-Hexene	Not Available	500	0.27	ND	D1	0.54
c-2-Pentene	Not Available	2,600	0.25	ND	D1	0.5
Carbon Tetrachloride	97,000	20	0.27	0.32	J,D1	0.54
Chlorobenzene (phenyl chloride)	210	100	0.27	ND	D1	0.54
Chloroform (trichloromethane)	85,000	20	0.21	0.13	J,D1	0.42
Cyclohexane	420	1,000	0.24	ND	D1	0.48
Cyclopentane	Not Available	1,200	0.27	0.13	J,D1	0.54
Cyclopentene	Not Available	2,900	0.2	ND	D1	0.4
Dichlorodifluoromethane	Not Available	10,000	0.2	0.5	L,D1	0.4
Ethane	180,000	Simple Asphyxiant*	0.5	6.1	D1,T	1
Ethylbenzene	170	20,000	0.27	ND	D1	0.54
Ethylene	270,000	500,000	0.5	0.74	J,D1,T	1
Isobutane	2,040	8,000	0.23	0.87	L,D1	0.46
Isopentane (2-methylbutane)	1,300	1,200	0.27	0.67	L,D1	0.54

Lab Sample ID	1104052-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
Isoprene	5	20	0.27	ND	D1	0.54
Isopropylbenzene (cumene)	100	500	0.24	ND	D1	0.48
m & p-Xylene (as mixed isomers)	80	1,700	0.27	0.43	J,D1	0.54
m-Diethylbenzene	70	460	0.27	ND	D1	0.54
Methyl Chloride (chloromethane)	Not Available	500	0.2	0.82	L,D1	0.4
Methylcyclohexane	150	4,000	0.26	ND	D1	0.52
Methylcyclopentane	1,700	750	0.27	ND	D1	0.54
Methylene Chloride (dichloromethane)	160,000	3,500	0.14	ND	D1	0.28
m-Ethyltoluene	18	250	0.11	ND	D1	0.22
n-Butane	1,200,000	8,000	0.2	1.4	L,D1	0.4
n-Decane	620	1,750	0.27	ND	D1	0.54
n-Heptane	670	850	0.25	ND	D1	0.5
n-Hexane	1,500	1,800	0.2	ND	D1	0.4
n-Nonane	2,200	2,000	0.22	ND	D1	0.44
n-Octane	1,700	750	0.19	ND	D1	0.38
n-Pentane	1,400	1,200	0.27	ND	D1	0.54
n-Propylbenzene	3.8	250	0.27	ND	D1	0.54
n-Undecane	Not Available	550	0.27	ND	D1	0.54
o-Ethyltoluene	Not Available	250	0.13	ND	D1	0.26
o-Xylene	380	1,700	0.27	ND	D1	0.54
p-Diethylbenzene	0.39	460	0.27	0.14	J,D1	0.54
p-Ethyltoluene	8.3	250	0.16	ND	D1	0.32
Propane	1,500,000	Simple Asphyxiant*	0.5	2.5	D1,T	1
Propylene	13,000	Simple Asphyxiant*	0.5	0.37	J,D1,T	1
Styrene	25	5,100	0.27	ND	D1	0.54
t-1,3-Dichloropropylene	Not Available	10	0.2	ND	D1	0.4

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Lab Sample ID	1104052-001					
Compound	Odor AMCV (ppb _v)	Short-Term Health AMCV (ppb _v)	LOD (ppb _v)	Concentrations (ppb _v)	Flags	SDL (ppb _v)
t-2-Butene	2,100	15,000	0.18	ND	D1	0.36
t-2-Hexene	Not Available	500	0.27	ND	D1	0.54
t-2-Pentene	Not Available	2,600	0.27	ND	D1	0.54
Tetrachloroethylene	770	1,000	0.24	0.12	J,D1	0.48
Toluene	170	4,000	0.27	0.25	J,D1	0.54
Trichloroethylene	3,900	100	0.29	ND	D1	0.58
Trichlorofluoromethane	5,000	10,000	0.29	0.36	J,D1	0.58
Vinyl Chloride	Not Available	26,000	0.17	ND	D1	0.34

^{*}A simple asphyxiant displaces air, lowering the partial pressure of oxygen and causing hypoxia at sufficiently high concentrations. ppbv - parts per billion by volume

ND - Not Detected.

NQ - Concentration can not be quantified.

LOD - Limit of Detection.

SDL - Sample Detection Limit (LOD adjusted for dilutions).

INV - Invalid.

- J Reported concentration is below SDL.
- L Reported concentration is at or above the SDL and is below the lower limit of quantitation.
- E Reported concentration exceeds the upper limit of instrument calibration.
- M Result modified from previous result.
- T Data was not confirmed by a confirmational analysis. Data is tentatively identified.
- D1 Sample concentration was calculated using a dilution factor of 4.02.

Table 4. TCEQ Long-Term Air Monitoring Comparison Values (AMCVs)

Please Note: The long-term AMCVs are provided for informational purposes only because it is scientifically inappropriate to compare short-term monitored values to the long-term AMCV.

Compound	Long-Term Health AMCV (ppb _v)	Compound	Long-Term Health AMCV (ppb _v)	
1,1,1-Trichloroethane	940	Cyclopentane	120	
1,1,2,2-Tetrachloroethane	1	Cyclopentene	290	
1,1,2-Trichloroethane	10	Dichlorodifluoromethane	1,000	
1,1-Dichloroethane	100	Ethane	Simple Asphyxiant*	
1,1-Dichloroethylene	86	Ethylbenzene	450	
1,2,3-Trimethylbenzene	25	Ethylene**	5,300	
1,2,4-Trimethylbenzene	25	Isobutane	800	
1,2-Dibromoethane	0.05	Isopentane (2-methylbutane)	120	
1,2-Dichloroethane	1	Isoprene	2	
1,2-Dichloropropane	10	Isopropylbenzene (cumene)	50	
1,3,5-Trimethylbenzene	25	m & p-Xylene (as mixed isomers)	140	
1,3-Butadiene	9.1	m-Diethylbenzene	46	
1-Butene	Not Available	Methyl Chloride (chloromethane)	50	
1-Pentene	Not Available	Methylcyclohexane	400	
2,2,4-Trimethylpentane	75	Methylcyclopentane	75	
2,2-Dimethylbutane (Neohexane)	100	Methylene Chloride (dichloromethane)	100	
2,3,4-Trimethylpentane	75	m-Ethyltoluene	25	
2,3-Dimethylbutane	99	n-Butane	800	
2,3-Dimethylpentane	85	n-Decane	175	
2,4-Dimethylpentane	85	n-Heptane	85	
2-Chloropentane (as chloroethane)	19	n-Hexane	190	
2-Methyl-1-Pentene +1-Hexene	50	n-Nonane	200	

Compound	Long-Term Health AMCV (ppb _v)	Compound	Long-Term Health AMCV (ppb _v)	
2-Methyl-2-Butene	50	n-Octane	75	
2-Methylheptane	75	n-Pentane	120	
2-Methylhexane	75	n-Propylbenzene	25	
2-Methylpentane (Isohexane)	100	n-Undecane	55	
3-Methyl-1-Butene	800	o-Ethyltoluene	25	
3-Methylheptane	75	o-Xylene	140	
3-Methylhexane	75	p-Diethylbenzene	46	
3-Methylpentane	100	p-Ethyltoluene	25	
4-Methyl-1-Pentene (as hexene)	50	Propane	Simple Asphyxiant*	
Acetylene	2,500	Propylene	Simple Asphyxiant*	
Benzene	1.4	Styrene	110	
Bromomethane (methyl bromide)	3	t-1,3-Dichloropropylene	1	
c-1,3-Dichloropropylene	1	t-2-Butene	Not Available	
c-2-Butene	Not Available	t-2-Hexene	50	
c-2-Hexene	50	t-2-Pentene	Not Available	
c-2-Pentene	Not Available	Tetrachloroethylene***	3.8	
Carbon Tetrachloride	2	Toluene	1,100	
Chlorobenzene (phenyl chloride)	10	Trichloroethylene	10	
Chloroform (trichloromethane)	2	Trichlorofluoromethane	1,000	
Cyclohexane	100	Vinyl Chloride	0.45	

^{*}A simple asphyxiant displaces air, lowering the partial pressure of oxygen and causing hypoxia at sufficiently high concentrations.

^{**}Long-term vegetation AMCV for Ethylene is 30 ppb.

^{***}Long-term vegetation AMCV for Tetrachloroethylene is 12 ppb.